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RESEARCH ARTICLE

Uncovering the Differential Molecular Basis of Adaptive Diversity in Three *Echinochloa* Leaf Transcriptomes

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Data Availability Statement: All RNAseq raw reads are available from the NCBI GeneBank (accession number SRP041999). All chloroplast cDNA sequences are available from the NCBI GeneBank (accession numbers KR058322, KR058323, KR058324, KR058325, KR058326, KR058327).

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Abstract

Echinochloa is a major weed that grows almost everywhere in farmed land. This high prevalence results from its high adaptability to various water conditions, including upland and paddy fields, and its ability to grow in a wide range of climates, ranging from tropical to temperate regions. Three *Echinochloa crus-galli* accessions (EC-SNU1, EC-SNU2, and EC-SNU3) collected in Korea have shown diversity in their responses to flooding, with EC-SNU1 exhibiting the greatest growth among three accessions. In the search for molecular components underlying adaptive diversity among the three *Echinochloa crus-galli* accessions, we performed *de novo* assembly of leaf transcriptomes and investigated the pattern of differentially expressed genes (DEGs). Although the overall composition of the three leaf transcriptomes was well-conserved, the gene expression patterns of particular gene ontology (GO) categories were notably different among the three accessions. Under non-submergence growing conditions, five protein categories (serine/threonine kinase, leucine-rich repeat kinase, signaling-related, glycoprotein, and glycosidase) were significantly (FDR, $q < 0.05$) enriched in up-regulated DEGs from EC-SNU1. These up-regulated DEGs include major components of signal transduction pathways, such as receptor-like kinase (RLK) and calcium-dependent protein kinase (CDPK) genes, as well as previously known abiotic stress-responsive genes. Our results therefore suggest that diversified gene expression regulation of upstream signaling components conferred the molecular basis of adaptive diversity in *Echinochloa crus-galli*.

Introduction

The genus *Echinochloa* consists of approximately 50 species across tropical and warm temperate zones [1]. *Echinochloa* can grow in a broad range of habitats, and these plants are common weeds that have caused serious crop losses in rice and maize fields, leading to 32% to 99%

collection and analysis, decision to publish, or preparation of the manuscript.

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reductions in crop yields [2–6]. In this genus, *Echinochloa oryzicola*, *E. crus-galli* var. *crus-galli*, *E. crus-galli* var. *praticola* have been major weeds in Korea [7,8]. Morphologically, the three species are barely indistinguishable, except for several traits, such as the presence of seed awn and the pattern of leaf growth. For example, a typical *E. oryzicola* has a relatively short seed awn compared to the long seed awn in *E. crus-galli* var. *crus-galli*. However, *E. crus-galli* var. *praticola* has no awn [8]. Cytologically, *E. oryzicola* is known to be tetraploid ($2n = 4x = 36$), whereas *E. crus-galli* is hexaploid ($2n = 6x = 54$) [9]. In eco-physiological traits, the three varieties are quite different. *Echinochloa oryzicola* grows only in flooded rice paddies, whereas *E. crus-galli* var. *praticola* mostly inhabits dry land and *E. crus-galli* var. *crus-galli* populates the areas from dry land to paddy rice fields [10]. Submergence tolerance of the three species has been systematically investigated using controlled environments [7]. In those studies, *E. oryzicola* was demonstrated to be the most submergence tolerant of the three species: The ranking for plant growth after 30 days of full (20 cm) submergence at the third leaf stage was *E. oryzicola* > *E. crus-galli* var. *crus-galli* > *E. crus-galli* var. *praticola*, supporting previous studies in the field [7].

One way to survive submergence stress is to change gene expression regulators, such as transcription factors. In submergence-tolerant species, certain transcription factors, such as the ethylene-responsive factor (ERF), have been demonstrated to play important roles in coping with submergence stress [11–14]. Under submergence (low O_2 concentration), the ethylene content in plants increases and can either stimulate or repress the elongation of internodes through ERF induction [12–16], such as *SNORKEL1* (*SK1*) / *SNORKEL2* (*SK2*) [17] and *Submergence1* (*Sub1*) [18,19]. Another way to cope with abiotic environmental stimuli is through activating signal transduction pathways, which generate physiological responses for adaptation. Although there are multiple types of external stimuli, the signal transduction shares common features. The generic components of the signal transduction pathway for plant responses to abiotic stress include (1) a receptor kinase for signal perception, (2) second messengers for signal amplification, (3) an array of signal transduction components, and (4) transcription factors (TF, involved in stress-responsive gene expression) [20]. Transducers and TFs involved in signal transduction pathways that respond to cold [21–26], salinity [27–30], and drought [31–33] have been well-studied. On the contrary, the signal perception components have not been well-defined, as is the case for receptor-like kinases (RLKs), which have been known for their roles in disease resistance [34], and in developmental process [35]. Recently, the role of RLKs in abiotic stress tolerance has been reported in several plant species [36–42], but many RLKs in plants were still functionally unknown [43–45]. Global gene expression studies of RLKs in rice using microarray after stress treatment revealed that certain genes encoding RLKs are constitutively expressed in various tissues, suggesting that they have important roles in the regulation of drought, salt, and cold responses [43]. Therefore, RLKs may play important roles in abiotic stress signaling, in addition to their roles in development and biotic stress.

The presence of differential eco-physiology in the closely related species or within the same species would be a useful system to study adaptive variation. Presence of *Echinochloa* accessions in Korea within the same species that exhibit adaptive variation would serve such a purpose. In this work, we adopted the three accessions of *E. crus-galli* which exhibit adaptive variation to explore the molecular basis of such difference. Because genomic resources for *Echinochloa* species have been limited so far, largely due to its large genome size and polyploidy ($2n = 4x = 36$ or $2n = 6x = 54$) [8], we used *de novo* transcriptome assembly as it has been an efficient approach for the large sized genomes of many crop species, saving cost and labor [46]. This approach was applied to two species of *Echinochloa* collected in China [47]. In this study, we investigated the composition and expression pattern of the three transcriptomes, in association with their ecological adaptive diversity. To identify major contributors, such as ERFs or

RLKs, which are responsible for abiotic stress adaption, we tested for differential gene expression patterns and significant enrichment of protein categories in their leaf transcriptomes. Our results will serve as basis for differential adaptive diversity in *Echinochloa* and related grass species.

Materials and Methods

Ethics statement

No specific permits were required for the described field study. No specific permissions were required for this location and activities. The location is not privately owned or protected in any way. The field studies did not involve endangered or protected species.

Plant materials and growth

Plants from three *E. crus-galli* accessions, EC-SNU1, EC-SNU2, and EC-SNU3, were collected in Seoul National University Experimental Farm Station in Suwon, Korea (E126°59'27.46", N37°16'12.05") in 2005. All *E. crus-galli* accessions were self-pollinated, and the progeny was processed for purification. One hundred seeds harvested from a single plant of each of the three *E. crus-galli* accessions that were selected based on parental characters for each accession, such as shape of seeds, existence of an awn and length of the outer glume (S1 Fig) [8]. The seeds were sown in petri dishes with wet filter paper and placed in a growth chamber (Hanbaek Scientific Ltd., Korea) under 25/20°C (day/night) and a 16 h photoperiod until the first leaf emergence. After the first leaf stage, twenty healthy seedlings of each *E. crus-galli* accession were selected and transplanted into to 11 cm diameter soil pots. Each *E. crus-galli* accession was isolated spatially to prevent contamination and allow it to grow to maturity. One plant per accession was selected as a typical morphological representative and its seeds were harvested. The seeds were stored at 4°C for 4 months to break their dormancy. Single-seed-descent lines were obtained by germinating seeds from each plant/species in Petri dishes and transplanting seedlings at the first leaf stage into pots filled with paddy soil; plants were grown in the glass house at the experimental farm station at Seoul National University, Suwon, Korea.

Submergence treatment and growth measurement

Three accessions of *E. crus-galli* seeds were water-soaked in petri dishes in a growth chamber at a temperature of 30/25°C and a 16/8 h photoperiod (day/night) until the first leaf emerged. Germinated seedlings were transplanted in pots filled with well-puddled paddy soil in the glass house at the Experimental Farm Station of Seoul National University at Suwon, Korea. Plants at the third leaf stage were placed in acrylic boxes under three flooding depths or 0 cm (water-logging, control) or 20 cm (complete submergence) for up to 30 days. For gross morphology analysis, the measurements of plant height and photographic analyses were performed at 7 days after flooding. The height was examined from the top of the soil to the top of the main plant stem. The dry weight was investigated at 30 days after flooding. The plants were removed from the soil, dried in an oven set at 80°C for 48 hours and weighed on a scale. For plant height and dry weight, statistical analyses were performed using the IBM SPSS 21.0 software for Windows. Data were analyzed with analysis of variance (ANOVA) and the means were considered to be significantly different if the Tukey-HSD post-hoc test showed $p < 0.05$.

Cytological analyses

For DNA content measurement, we performed flow cytometric analysis of the three *E. crus-galli* accessions using one sorghum accession (*Sorghum bicolor* cv. Pioneer 8695) as an internal

standard, and one *E. oryzicola* accession as a tetraploid control. Leaf tissues ($\sim 0.5 \text{ cm}^2$) were chopped with a double-edge razor in a petri dish containing 0.5 mL nuclear extraction buffer (CyStain UV precise P/Solution A, PARTEC, Germany). After samples were chopped, they were filtered through a 30 μm mesh filter. After filtration, 2.0 mL of a staining solution containing 4',6'-diamidino-2-phenylindole (DAPI) fluorochrome (CyStain UV precise P/Solution B, PARTEC, Germany) was added. Flow cytometry was performed using Ploidy Analyzer (CyFlow PA, PARTEC, Germany). Fluorescence peaks were estimated using the WinMDI software package (version 2.9, <http://en.bio-soft.net/other/WinMDI.html>). Three replications per leaf were analyzed.

For chromosome counting, young root tips ($\sim 0.5 \text{ cm}$) were sampled and placed into 1.5 mL microtubes filled with 0.002 M 8-hydroxyquinolin. After 2 hours, the solutions were removed 3:1 absolute alcohol:glacial acetic acid was added at room temperature for one day. The fixed samples were washed with distilled water for 2 minutes, hydrolyzed with 5N HCl for 45 minutes and the root tips were rinsed with distilled water for 2 min. The samples were stained using Schiff's stain (0.01 M pararosaniline hydrochloride, 0.01 M Potassium disulfite, 0.15N HCl) for 2 hours in darkness. After staining, samples were washed with distilled water for 10 minutes. The root tips were soaked in enzyme solution (cellulysin and macerace in 10^{-3}M EDTA) for 45 minutes, placed on a slide glass and tapped with a stainless steel stick. The samples were observed under 1,000X magnification using a Photo Microscope (Axiophot, ZEISS, Germany).

Phylogenetic analysis

matK and *rbcL* sequences from six species (*E. crus-galli* EC-SNU1, EC-SNU2, and EC-SNU3 accessions, *E. oryzicola*, *Oryza sativa*, and *Sorghum bicolor*) were used for phylogenetic tree construction and estimation of divergence time. The *matK* and *rbcL* sequences of three *E. crus-galli* accessions were retrieved from the previous chloroplast genome sequencing of three *E. crus-galli* accessions (Nah et al., unpublished), whereas the sequences of *E. oryzicola* (GenBank ID: KJ000048) [48], *O. sativa* (GenBank ID: NC_001320.1) and *S. bicolor* (GenBank ID: NC_008602.1) were obtained from GenBank (<http://www.ncbi.nlm.nih.gov/gene>). Multiple sequence alignment of *matK* cDNA sequences was performed using Clustal W (<http://www.clustal.org/clustal2/>). After sequence alignment, a phylogenetic tree was constructed using the BEAST package (<http://beast2.org/>) with a Hasegawa-Kishino-Yano (HKY) model of evolution used for Bayesian MCMC analysis. The tree was displayed through FigTree (version 1.4.2, <http://beast.bio.ed.ac.uk/figtree>). *matK* and *rbcL* sequences from the three *E. crus-galli* accessions were deposited into NCBI GenBank (EC-SNU1_ *rbcL*: KR058322; EC-SNU2_ *rbcL*: KR058323; EC-SNU3_ *rbcL*: KR058324; EC-SNU1_ *matK*: KR058325; EC-SNU2_ *matK*: KR058326; EC-SNU3_ *matK*: KR058327).

RNA isolation, cDNA library construction, and sequencing

Total RNA was extracted from the fourth leaves of three *E. crus-galli* accessions before the first tillering, using Trizol reagents (Invitrogen, USA) according to the manufacturer's instructions. Quantity and quality of total RNA were analyzed using a Nanodrop 2000 spectrophotometer (Nanodrop, USA) and gel electrophoresis (Takara, Japan), respectively. cDNA library construction was performed using the TruSeq kit (Illumina, USA), at the National Instrumentation Center for Environment Management (NICEM), at Seoul National University, Korea. Sequencing was performed by 101bpX2 (paired-end sequencing) for Illumina HiSeq2500 (Illumina, CA, USA) at NICEM. All reads were deposited in the National Center for Biotechnology Information (NCBI) center under the SRA accession of SRP041999.

De novo assembly, annotation, and gene ontology categorization

For *de novo* contig assembly, a total of 78,500,009 paired end reads were obtained in three libraries; the EC-SNU1 library has 22,713,977 paired reads, the EC-SNU2 library has 29,405,121 paired reads, and the EC-SNU3 library has 26,380,911 paired reads. We filtered the raw reads in two steps: the low quality reads were removed by the trimming script from the FASTX toolkit (version 0.0.12; http://hannonlab.cshl.edu/fastx_toolkit/). Reads with q-values less than 30 were trimmed, and then PCR duplicates were removed by the trimming script from the NGS QC toolkit (version 2.3; <http://59.163.192.90:8080/ngsqctoolkit/>). We used Trinity (r2013-02-25; <http://trinityrnaseq.sourceforge.net/>) for *de novo* assembly of the *E. crus-galli* leaf transcriptome reads (min_kmer_cov = 6, JM = 20G, CPU = 2, min_contig_size = 300). After assembly, we performed CD-hit (<http://www.bioinformatics.org/cd-hit/>) with an identity of 95% to remove redundancy. Next, we filtered out known transposable elements from the assembled contigs using mipsREdat_9.3p_ALL.fasta (<http://www.mmnt.net/db/0/0/ftp.mips.embnet.org/plants/REdat>) using BLASTN with an e-value less than $1E^{-10}$, an alignment identity greater than 90%, and an alignment length greater than 100bp. Annotation was performed using BLASTX with an e-value less than $1E^{-5}$ against *Oryza sativa* peptide sequence data with annotation information from Rice Genome Annotation Project (ftp://ftp.plantbiology.msu.edu/pub/data/Eukaryotic_Projects/o_sativa/annotation_dbs/pseudomolecules/version_7.0/all.dir/all.pep). For gene ontology (GO) categorization, we used DAVID (<http://david.abcc.ncifcrf.gov/>), followed by calculation of relative ratios. The relative ratio was estimated using the number of contigs assigned to an individual GO category divided by the total number of contigs used for GO assignment for each accession.

Identification of differentially expressed genes (DEG), hierarchical clustering, and GO-enrichment assay

Using assembled contigs from EC-SNU1 as a reference, we mapped raw reads from SRP041999 to quantify the expression level for DEG identification. For this purpose, run_DE_analysis.pl from edgeR (with $q < 0.05$) implemented in Trinityrnaseq_r2013-02-25 package was used. This script identified DEGs in pair-wise comparisons using mapped reads. Next, we calculated the FPKM (Fragments per Kilo base-pair per Million mapped fragments) value for individual genes in EC-SNU1, EC-SNU2, and EC-SNU3 using run_RSEM_align_n_estimate.pl from the Trinityrnaseq_r2013-02-25 package. FPKMs, when found with a value greater than 2 in at least one of the accessions, were used for gene expression clustering. For gene expression clustering, we used Pearson's correlation with average distance at GENE-E (<http://www.broadinstitute.org/cancer/software/GENE-E/download.html>) using the \log_2 value of FPKM. For the GO enrichment assay, we used DAVID (<http://david.abcc.ncifcrf.gov/>). We first selected contigs with significant matches to the Arabidopsis protein database (TAIR10) and 18,596 contigs matched with TAIR10 (e-value $< 1E^{-10}$) were used as a background gene set for DAVID. Contigs from DEG were converted to TAIR10 protein IDs based on their homologies and run as a sample gene set for DAVID. To select significant GO categories, we applied the q-value approach using a False Discovery Rate (FDR) threshold of less than 0.05. All other runs, such as BLASTN and BLASTX were performed using the LINUX operating system (CentOS, <http://www.centos.org/download/>).

Quantitative real time PCR verification

For qRT-PCR analysis, leaf samples were collected from the full growth of 4th leaf stage. Total RNA was extracted from 100 mg of leaf tissue using the RNeasy Plant Mini Kit (Qiagen, CA,

USA). Single-stranded cDNA was synthesized from 500 ng of total RNA using TOP-script Reverse Transcriptase (Enzynomics, Korea). *eEF-1 α* was used as a housekeeping gene to normalize the three *E. crus-galli* accessions. Primers for 19 genes (2 Serine/threonine receptor kinases, 4 Leucine-rich repeat receptor kinases, 5 Cysteine-rich protein kinases, and 8 Calcinurin B-like kinases) were used (S1 Table). qRT-PCR reactions were carried out using TOP-realTM qPCR 2X PreMIX for SYBR Green (Enzynomics, Korea) with a final volume of 20 μ l in a Rotor-Gene Q 2plex real-time PCR machine (Qiagen, CA, US). The qRT-PCR cycle program was as follows: (1) 95°C for 15 min for initial denaturation, (2) 95°C for 10 sec for denaturation, (3) 55°C for 15 sec for annealing, (4) 72°C for 20 sec for elongation, and then the cycles from (2)-(4) were repeated 45 times. Relative quantification of each single gene expression was evaluated using the delta-delta Ct method [49].

Results

Morphological, cytological, and physiological overview of three *E. crus-galli* accessions

We investigated seed morphology of three *E. crus-galli* accessions, particularly, seed size and awn (S1 Fig). EC-SNU1 resembled more to the *E. oryzicola* than to *E. crus-galli*, but still exhibited relatively smaller size and longer awn than those of *E. oryzicola*. The seed shape of EC-SNU1 resembled the typical C-type *E. oryzicola* [50], but cytological analysis below revealed hexaploidy nature (S2 and S3 Figs), a major characteristic of *E. crus-galli*. EC-SNU2 also exhibited smaller seed size and the longest awn among all accessions, which are typical traits of *E. crus-galli* var. *crus-galli*. Finally, EC-SNU3 displayed in smallest seed size without awn, the typical traits of *E. crus-galli* var. *pratensis*.

To obtain basic genetic information on these three *Echinochloa* accessions, we investigated their ploidy levels. Flow cytometry measurements showed that all three accessions carry DNA content (ANOVA, $p < 0.05$) similar to hexaploids (S2 Fig). Chromosome counting finally confirmed that they are hexaploids ($2n = 6x = 54$) (S3 Fig). This property is a major characteristic of *E. crus-galli*, which distinguishes it clearly from the tetraploid *E. oryzicola* (S2 and S3 Figs).

To validate the eco-physiological implications of the fact that EC-SNU1 and EC-SNU2 were collected from paddy rice field, whereas EC-SNU3 was collected from dry land, we investigated the physiological responses for the three accessions under controlled submergence conditions, and evaluated their growth by measuring their leaf growth and biomass yield. Compared to control (0 cm) and the full submergence (20 cm) treatment gave significant differences in growth among the three accessions, particularly EC-SNU1 exhibited significantly ($p < 0.05$) more growth compared to the other two accessions (S4 Fig).

De novo assembly of three *E. crus-galli* leaf transcriptomes

To investigate the genomic constituent of the three *E. crus-galli* accessions, we performed *de novo* assembly of raw reads from three *E. crus-galli* leaf transcriptomes generated by Illumina HiSeq. Initially, we obtained 18,368,735 reads from EC-SNU1 (81% recovery ratio), 23,428,201 reads from EC-SNU2 (80% recovery ratio), and 21,122,046 reads from EC-SNU3 (80% recovery ratio). Using Trinity with 25-mers, as recommended by the published procedure [51], we obtained 18,368,735 reads from EC-SNU1 (81% recovery ratio), 23,428,201 reads from EC-SNU2 (80% recovery ratio), and 21,122,046 reads from EC-SNU3 (80% recovery ratio). As a result, we retrieved 31,160 contigs with N50 of 1,435 bp from EC-SNU1, 28,814 contigs with N50 of 1,278 bp from EC-SNU2, and 30,048 contigs with N50 of 1,411 bp from EC-SNU3 (Table 1). Because we used cDNA fragments sequenced by paired-end (101bpX2) reading with

Table 1. Statistics of *de novo* assembly of *Echinochloa* leaf transcriptomes.

	EC-SNU1	EC-SNU2	EC-SNU3
Total number of contigs	31,023	28,882	30,048
Total bases (bp)	32,586,794	31,527,271	31,527,271
Minimum contig length (bp)	301	301	301
Maximum contig length (bp)	12,060	7,412	11,788
Average contig length (bp)	1,050	977	1,049
N ₅₀ length (bp)	1,435	1,278	1,411

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a minimum of 100 bp between the two ends, 300 bp was chosen as the minimum contig size. As a result, the average length of a contig was 1,050 bp, 976bp and 1,049bp in EC-SNU1, EC-SNU2, and EC-SNU3, respectively (Table 1). The range of contig sizes showed that the majority of contigs were 300–1,500bp, covering 78.2% (24,361 contigs) of the EC-SNU1 library, 81.8% (23,565 contigs) of the EC-SNU2 library, and 78.5% (23,585 contigs) of the EC-SNU3 library (Fig 1A).

Annotation and gene ontology categorization of three *E. crus-galli* leaf transcriptomes

For assembly validation and annotation, BLASTX against known protein databases of *Oryza sativa*, *Sorghum bicolor*, and *Setaria italica* was performed. Using a threshold e-value of less than E^{-05} and an aligned amino acid length of 50aa, 63–67% of the translated *E. crus-galli* contigs were matched against the *Oryza sativa* protein database, 65–69% against the *Sorghum bicolor* protein database, and 65–70% against *Setaria italica* protein database (Fig 1B). In all three accessions, the *C₄* grass *Setaria italica* showed the highest match among three grass species protein databases. Next, the annotated contigs were subjected to gene ontology (GO) categorization for three categories (biological, cellular, and molecular process), and the three accessions showed similar relative gene ratios of genes in each individual category, and exhibited the same categories with similar ratios of contigs in the leaf transcriptomes (S5 Fig, S2, S3, and S4 Tables). The top five categories were also well-conserved among the three accessions: For biological process, the top five categories included biological process (12.7–13.6% of an individual transcriptome), cellular process (12.0–12.2%), metabolic process (11.0–11.3%), biosynthetic process (7.5–7.9%), and response to stress (4.7–4.9%). For cellular process, cellular component (13.8–14.6%), plastid (13.0–13.7%), membrane (11.7–12.4%), plasma membrane (8.6–8.9%), and nucleus (8.4–8.6%) were included in the top five. For molecular process, molecular function (17.2–18.5%), catalytic activity (10.8–11.1%), protein binding (10.2–10.8%), hydrolase activity (8.5–8.7%), and DNA binding (8.0–8.8%) were included in the top five. Overall, GO categories and their relative ratios were well conserved among the three leaf transcriptomes. To investigate the homolog ratios shared by three leaf transcriptomes, we performed pair-wise BLASTN with an e-value of less than $1E^{-10}$. The majority of the transcriptome consisted of homologs, ranging from 80.4% to 92.2% (S6 Fig).

Differential gene expression of three *E. crus-galli* leaf transcriptomes

Although EC-SNU1 and EC-SNU2 showed no significant physiological and growth difference in submergence response (S4 Fig), it is possible that they are different at molecular level in relation to other abiotic stress. To perform systematic investigation of overall abiotic stress-related gene expression in three accessions, we conducted pair-wise comparisons of the DEGs and detected the presence of significant ($q < 0.05$) differential gene expression. The pattern of DEG

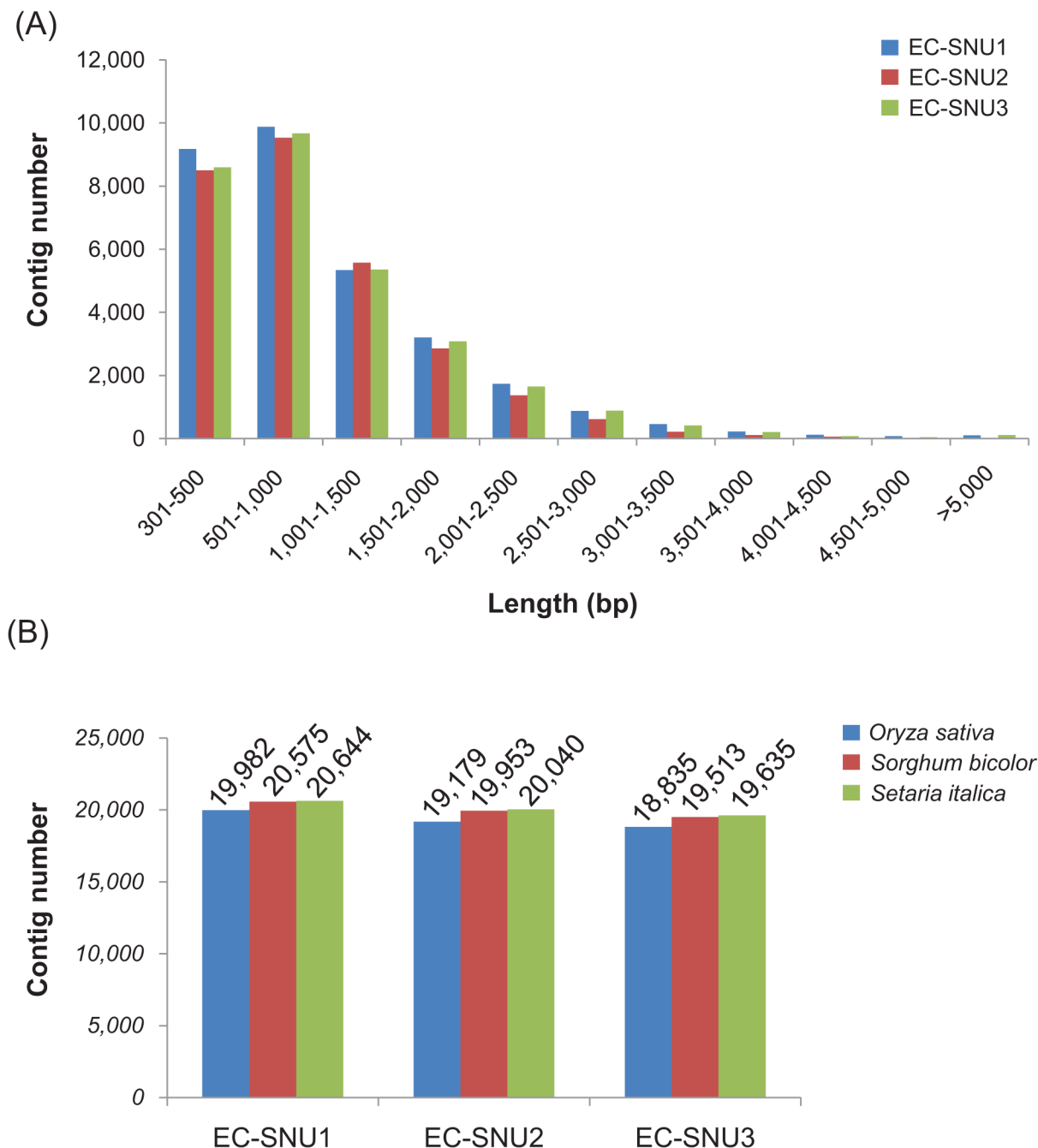


Fig 1. (A) Contig size distribution of *de novo* assembly of transcriptomes from three *Echinochloa* accessions. (B) Assembly validation of known grass protein databases for rice (*Oryza sativa*), sorghum (*Sorghum bicolor*), and foxtail (*Setaria italica*).

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distribution exhibited a bias toward up-regulation in the EC-SNU1 contigs in all pair-wise comparisons (Fig 2A and 2B). However, the comparisons for EC-SNU2 and EC-SNU3 exhibited a relatively similar number of contigs in the DEG distributions (Fig 2C). We counted the total number of contigs belonging to these DEGs. In each pair-wise comparison, 1,061 contigs were up-regulated in EC-SNU1 compared to 399 up-regulated in EC-SNU2, 936 were

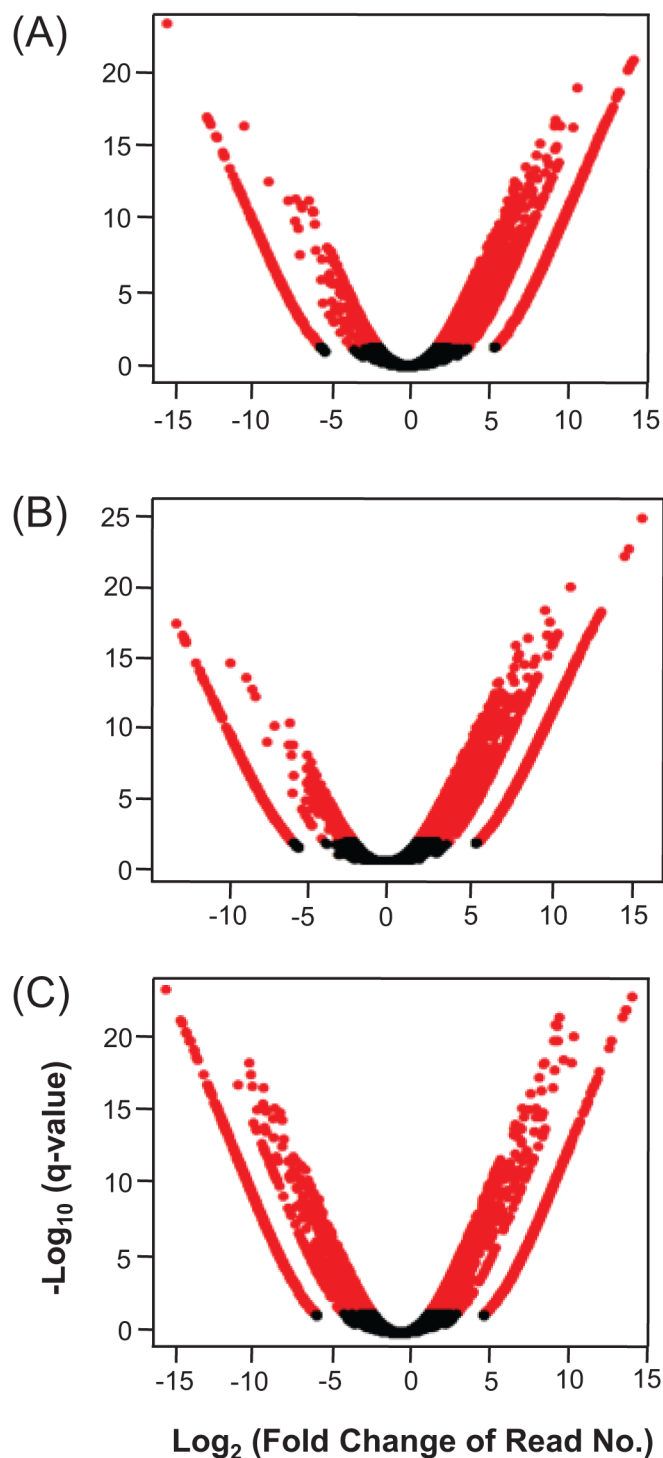


Fig 2. Distribution of \log_2 fold change of DEGs in pair-wise comparison; (A) EC-SNU1 vs. EC-SNU2, (B) EC-ENU1 vs. EC-ENU3, and (C) EC-SNU2 vs. EC-SNU3.

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up-regulated in EC-SNU1 compared to 269 up-regulated in EC-SNU3, and 857 were up-regulated in EC-SNU2 compared to 756 up-regulated in EC-SNU3 ([S7 Fig](#)).

To examine whether a large number of up-regulated DEGs are enriched with particular protein categories, GO enrichment analysis was performed. Five groups of representative enriched protein categories from SWISS-PROT and the Protein Information Resource Database were retrieved: (1) serine/threonine protein kinase (STK) (GO:0004674), (2) leucine-rich repeat receptor-like kinase (LRR-RLK) (GO:0035872), (3) signaling-related protein (GO:0023052), (4) glycoprotein (GO:0047965), and (5) glycosidase (GO:0016798) (Fig 3A). A hierarchical clustering diagram of STK and LRR-RLK groups of annotated transcripts were significantly up-regulated in EC-SNU1 relative to both EC-SNU2 and EC-SNU3 (Fig 3B and 3C, S5 and S6 Tables). Although there was up-regulation of DEGs of EC-SNU2 and EC-SNU3 against those of EC-SNU1 in STK and LRR-RLK groups (Fig 3B and 3C), majority of up-regulation of DEGs was enriched in EC-SNU1. Of the RLKs, particularly Ca^{2+} -related signaling kinases were identified with dominant number in EC-SNU1 (S8A Fig, S7 Table), which are known participants in signal transduction. A hierarchical clustering diagram of signaling-related and glycoprotein groups of annotated transcripts were significantly up-regulated in both EC-SNU1 and EC-SNU2, relative to EC-SNU3 (Fig 3D and 3E, S8 and S9 Tables). Finally, a hierarchical clustering diagram of glycosidase group of annotated transcripts was significantly up-regulated in EC-SNU2, relative to EC-SNU1 and EC-SNU3 (Fig 3F, S10 Table). In addition to five GO categories, we observed the homologs of previously known abiotic stress-related genes in Arabidopsis were identified as DEGs and outnumbered in EC-SNU1, compared to EC-SNU2 and EC-SNU3 (S8B Fig, S11 Table).

We were particularly interested in RLK and 19 randomly selected DEGs were validated for relative expression using qRT-PCR (S1 Table), and of them, six genes were displayed (Fig 4). In most cases of RLK genes, the relative expression ratio was highest in EC-SNU1 (Fig 4).

We also investigated known submergence tolerance genes, *SNORKEL1/2* (*SK1/2*) [17] and *Submergence1A* (*Sub1A*) [18], in our transcriptomes based on sequence homology, but we could not detect any homologs of *Sub1A/B/C* or *SK1/2*. Instead, we observed several ERF homologs that were up-regulated in EC-SNU1 when EC-SNU1 was compared to EC-SNU3, and in EC-SNU2 when EC-SNU2 was compared to EC-SNU3. In both cases, the homolog of AT1G28370 (ERF domain 11), from the B-1 subfamily of the ERF/AP2 transcription factor family, was observed. An association between AT1G28370 and abiotic stress has been reported in previous studies [52–54].

Phylogenetic relationship among the three *E. crus-galli* accessions

The five enriched GO protein categories were detected in a step-wise manner in pair-wise comparisons, such that enrichment of up-regulated glycosidase category was found in EC-SNU2 against EC-SNU3 and EC-SNU1, enrichment of up-regulated of the signaling-related and glycoprotein categories were found in EC-SNU1 and EC-SNU2 against EC-SNU3, and finally, enrichment of up-regulated serine/threonine kinase and LRR kinase, categories were found in EC-SNU1 against EC-SNU2 and EC-SNU3 (Fig 3A). This observation raised the question as to whether such a step-wise acquisition supports our current understanding of the evolutionary divergence order. To answer this question, we established the phylogenetic relationship among the three accessions using two chloroplast genes, *matK* and *rbcL*. We used the *matK* and *rbcL* genes because they have been known as the most rapidly evolving coding regions in the chloroplast genome, and their sequence diversity provides a high level of discrimination to establish evolutionary relationships among angiosperm species [55]. Thus, these two genes have been selected to establish phylogenetic relationship by the Consortium for the Barcode of Life (http://www.barcoding.si.edu/plant_working_group.html). Evolutionary relationship from the phylogenetic trees using both *matK* and *rbcL*, suggested that EC-SNU1 diverged from the most

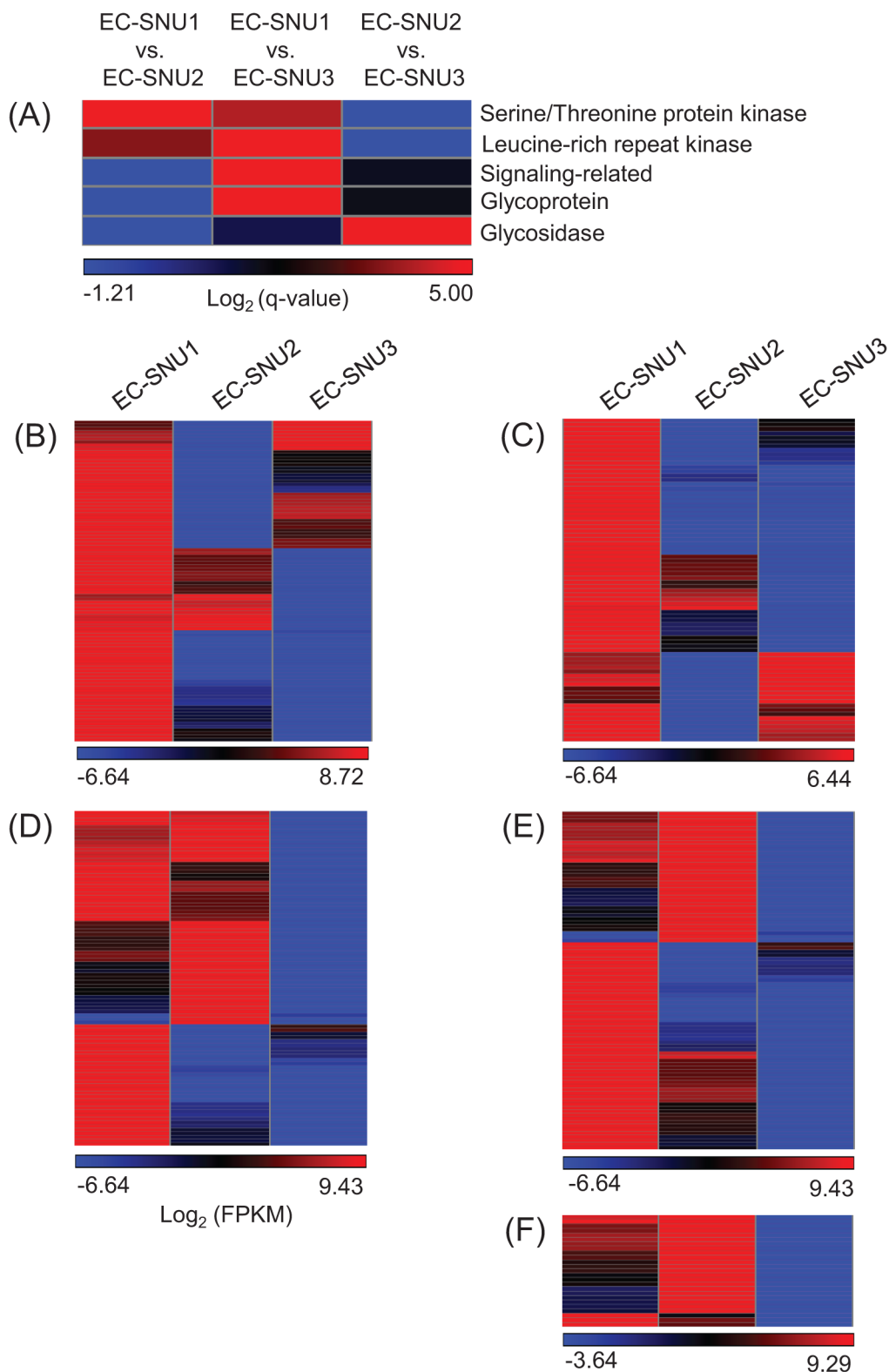


Fig 3. (A) Significant GO category enrichment in pair-wise comparisons. (B) Hierarchical clustering of DEG belonging to serine/threonine protein kinases. (C) Hierarchical clustering of DEG belonging to leucine-rich repeat kinases. (D) Hierarchical clustering of DEG belonging to signaling-related proteins. (E) Hierarchical clustering of DEG belonging to glycoprotein. (F) Hierarchical clustering of DEG belonging to glycosidase.

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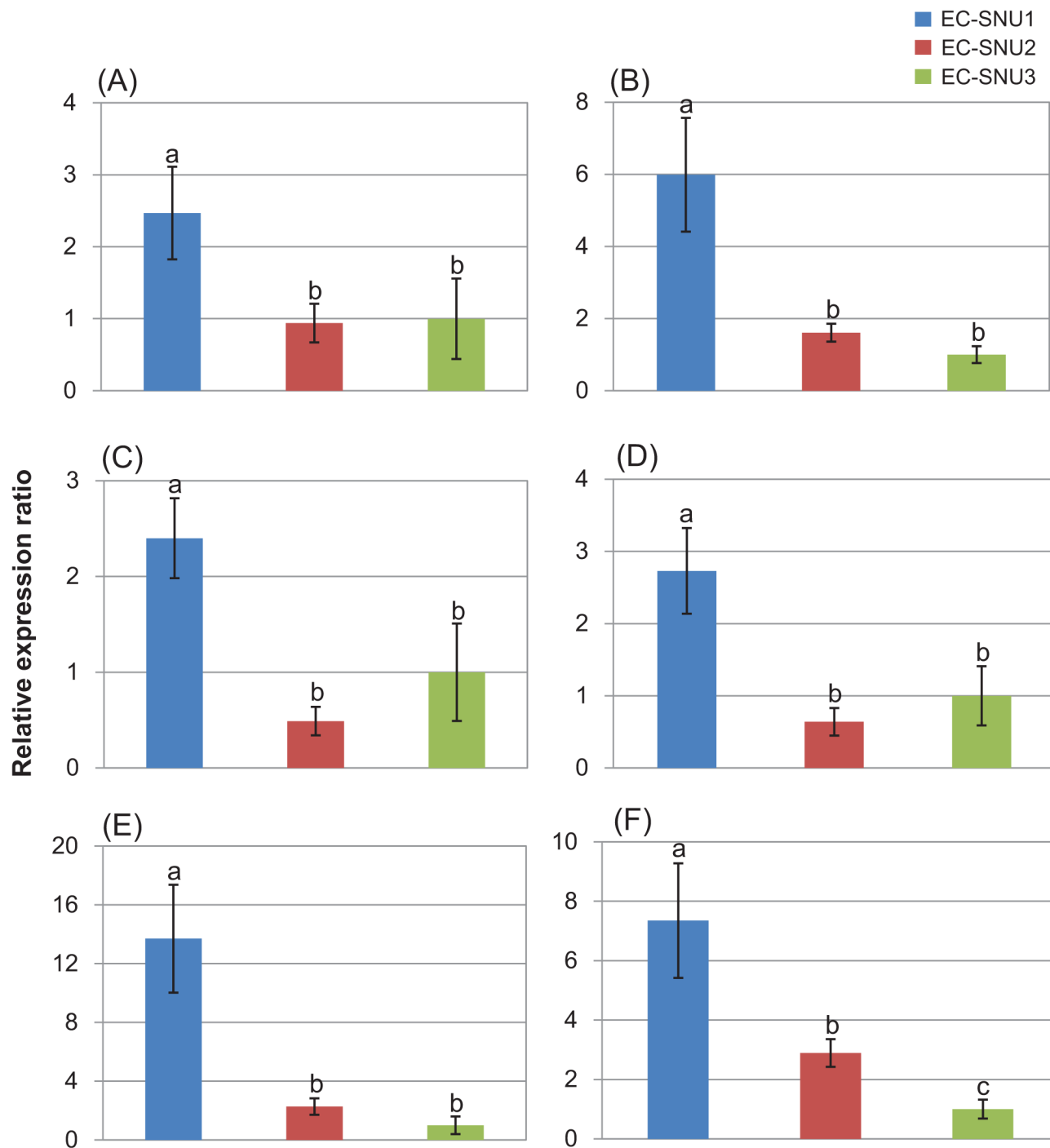


Fig 4. qRT-PCR validation of significantly up-regulated DEGs in EC-SNU1; (A) SOS3-interacting protein 1, (B) Calcium-dependent kinase 6, (C) Calmodulin 1, (D) LRR-RLK, (E) Cysteine-rich RLK 29, and (F) Cysteine-rich RLK 40.

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recent common ancestor of EC-SNU2 and EC-SNU3, followed by split into EC-SNU2 and EC-SNU3 (S9 Fig).

Discussion

Based on transcriptome sequence homology against protein databases using BLASTX, the C4 plant *Echinochloa* was best matched against another C4 species, foxtail millet (*Setaria italica*). Foxtail millet is a close relative of C4 bioenergy crops, such as switchgrass (*Panicum virgatum*), napiergrass (*Pennisetum purpureum*), and pearl millet (*Pennisetum glaucum*) [56], and its whole genome was sequenced [57], serving as a useful C4 reference system for other C4 grass species [58]. With foxtail millet, 60~70% of transcripts of *Echinochloa* were significantly matched, indicating the conservation of common genetic components among these C4 grasses. At the same time, the detection of 30~40% of difference in the transcriptome indicates that they might be mis-assembled contigs or *Echinochloa*-specific transcripts.

Although EC-SNU1 and EC-SNU2 showed no significant difference in flooding response based on flooding treatment experiment, they are still useful system together with EC-SNU3 in the comparative aspect, because they exhibited the variation at gene expression level in relation to abiotic stresses. Interestingly, we found that DEGs of EC-SNU1 significantly up-regulated, compared to those of EC-SNU2 and EC-SNU3, at gene expression level of certain abiotic-stress related GO categories, particularly RLK genes. To cope with unfavorable environmental changes, plants need to develop dynamic endogenous mechanisms, such as signal transduction. RLKs have been known to play important roles in sensing abiotic stress and signaling to the downstream network [59–62]. RLK forms a large gene family, consisting of over 600 genes in *Arabidopsis* [45,63] and over 1,000 genes in rice [45], indicating active and diverse roles in signaling from the environment stimuli. RLK-encoding genes are known to be increased by tandem duplication, and the presence of a large number of paralogs provides a source for genetic diversity [44,64,65]. Previous studies have demonstrated that RLKs act as key regulators in various abiotic stress responses in many plant species, such as *Arabidopsis thaliana* [66–69], *Oryza sativa* [41,70,71], *Medicago truncatula* [36], and *Glycine soja* [72]. These abiotic stress-related RLKs include leucine-rich repeat receptor-like protein kinases (LRR-RLK) [38,73], cysteine-rich RLK (CRK) [66], and calcium/calmodulin-dependent kinase (CaMK) [66], all of which were found as DEGs in our analysis. One important feature of RLKs is that they are upstream components in signal transduction. The significant up-regulation of RLK-encoding genes in EC-SNU1 reflected that this accession might have evolved in a different way within *E. crus-galli* via modification of regulatory regions. Although we tested only submergence, EC-SNU1 might have acquired additional abiotic stress tolerance traits, a hypothesis that needs further testing. This result suggests that evolutionary processes selected one or more upstream genes in the signaling pathway, probably because this conferred more efficient and dramatic impacts on the downstream pathways.

Moreover, our results suggested that expression pattern change might be an efficient strategy for plants to cope with abiotic stress, rather than generating new genes. Although it is possible that transcription of RLK-encoding genes can also be induced by external stimuli, constitutive up-regulation, as seen in the tolerant accession, can be a strategy for rapidly coping with adverse environmental conditions. For example, the gene expression profiling of 17 calcium-dependent protein kinase (CDPK) genes in grapevine exhibited constitutive expression that is prevalent in pollen [74], indicating that constitutive expression is one of the adaptive signaling strategies. Therefore, adaptive diversity can be achieved by inducing differential expression of signaling genes via modification of their regulatory regions. Moreover, previous studies indicated that modification of gene expression patterns can enhance abiotic tolerance. For example, the study on RLK showed that water-stress related tolerance was enhanced in *RPK1*-overexpressing plants, whereas loss-of-function mutants showed down-regulation of stress-related genes [68]. Additionally, rice *OsSIK1*-overexpressing plants exhibited enhanced salt

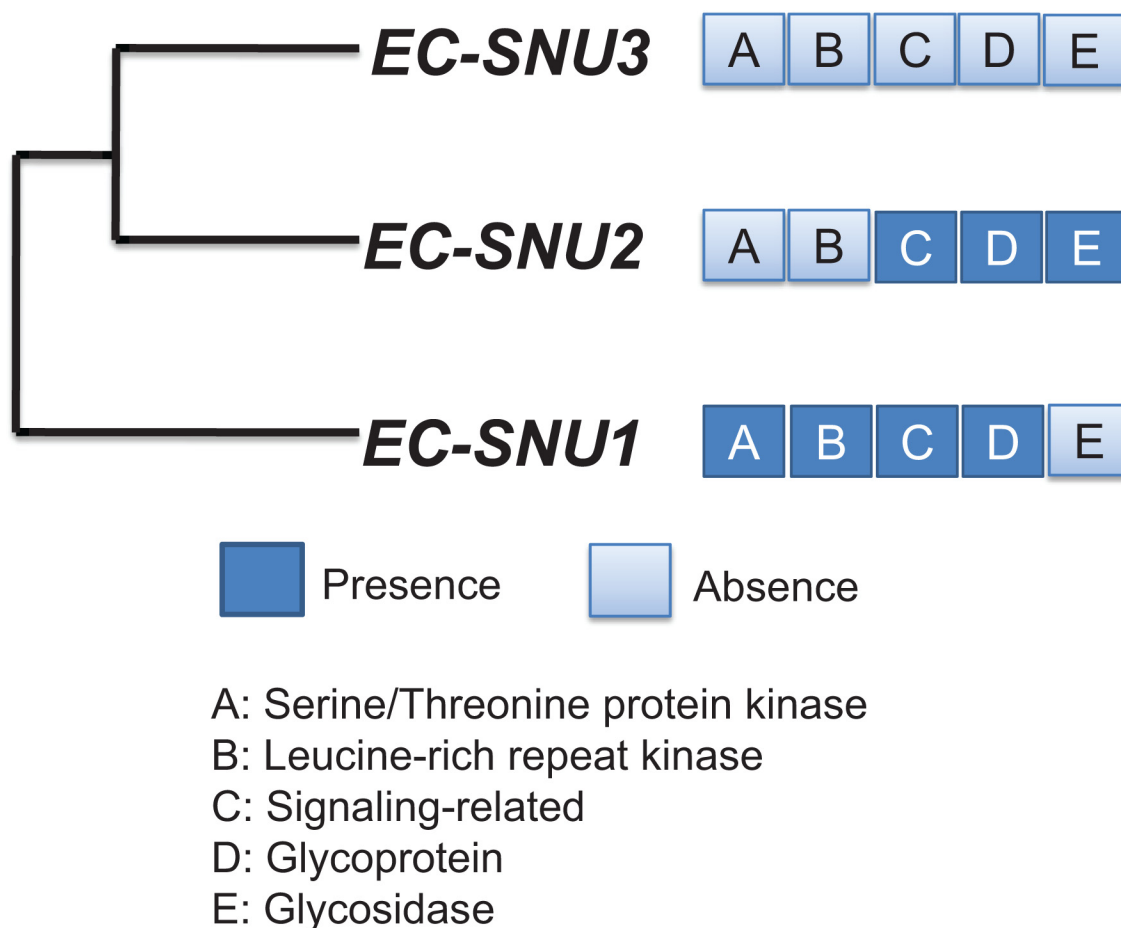


Fig 5. The step-wise change of gene expression pattern in the phylogenetic relationship of three *E. crus-galli* accessions.

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and drought tolerance, whereas loss of function of *OsSIK1* resulted in sensitivity to these stresses [37].

Another observation is that the step-wise loss and acquisition of gene expression groups corresponded to the phylogenetic relationship of the three accessions (S8 Fig). After divergence from EC-SNU1, the common ancestor of EC-SNU2 and EC-SNU3 appears to have lost gene expression up-regulation in two GO categories (STK and LRR-RLK), and then EC-SNU3 alone has lost up-regulation in two additional GO categories (signaling-related and glycoprotein). On the contrary, up-regulation of glycosidase appears to be acquired in EC-SNU2 after divergence from EC-SNU3. Based on this observation, we generated a model for the stepwise loss and acquisition of expression diversification in five GO categories during evolution (Fig 5). The availability of regulatory sequences for all three accessions will be helpful to validate this result by determining the diversification of the regulatory regions.

Unlike the RLKs, we were unable to detect orthologs of both *Sub1A/B/C* and *SK1/2* in three *E. crus-galli* leaf transcriptomes. The absence of *Sub1* or *SK* in *E. crus-galli* may be due to the limitations of the transcriptome size, or because *E. crus-galli* does not carry these genes. Instead, we identified one *ERF* homolog (AT1G28370) in the DEGs. This *E. crus-galli* *ERF* homolog might be the candidate for a submergence-responsive *ERF*. In addition, *ERF* is known to respond to drought, salt, and freezing stresses, indicating that *ERFs* might regulate multiple abiotic stresses [75]. Therefore, it may be useful to regulate multiple abiotic stress pathways

using a single gene, such as genes encoding ERF or RLK, to generate stress-tolerant plants. We also identified new groups of proteins, glycoprotein and glycosidase, which might be related to abiotic stress tolerance. Although a very few studies have been done, recent reports provided involvement of these groups of proteins in abiotic stress tolerance, by modifying cell wall metabolism [76–80].

Based on our analyses, we conclude that significantly up-regulated DEGs in EC-SNU1, including RLK-encoding genes, can be useful source for future studies of adaptive diversity, as well as for identification of novel abiotic stress-related genes in other grass species.

Conclusions

The three *E. crus-galli* leaf transcriptomes were generated and served as a useful system to identify and explore the molecular basis for environmental adaptability among accessions. Comparison of the three accessions revealed that family of RLK-encoding genes, which has been known to play important roles in signal transduction in both biotic and abiotic stresses, were major contributors to adaptive diversity. Moreover, previously known abiotic stress-related genes were also observed in these *Echinochloa* leaf transcriptomes. The fact that RLK-encoding genes and previously known abiotic stress-related genes were significantly up-regulated primarily in EC-SNU1 compared to EC-SNU2 or EC-SNU3 suggests the presence of variation in environmental adaptability, even within the *E. crus-galli* species. The phylogenetic relationship among the three accessions also implies that the regulatory changes in many RLK-encoding genes have been maintained in EC-SNU1, whereas some of them were lost in EC-SNU2 and EC-SNU3 during evolution. Our analysis therefore implies that gene expression modification of signaling components plays an important role in adaptive diversity in *Echinochloa* accessions.

Supporting Information

S1 Fig. Seed morphology of *Echinochloa* accessions. (A) *E. oryzicola*, (B) EC-SNU1, (C) EC-SNU2, and (D) EC-SNU3.
(PDF)

S2 Fig. Flow cytometric measurement of DNA content in *E. oryzicola* and three *E. crus-galli* accessions; (A) *E. oryzicola* (tetraploid control), (B) EC-SNU1, (C) EC-SNU2, and (D) EC-SNU3.
(PDF)

S3 Fig. Chromosomes from *E. oryzicola* and three *E. crus-galli* accessions; (A) *E. oryzicola* (tetraploid control), (B) EC-SNU1, (C) EC-SNU2, and (D) EC-SNU3.
(PDF)

S4 Fig. Relative plant height (A) and relative dry weight (B) of *Echinochloa* accessions at 7 days after submergence treatment.
(PDF)

S5 Fig. GO categorization of assembled contigs from three *Echinochloa* leaf transcriptomes; (A) Biological process, (B) Cellular process, and (C) Molecular process.
(PDF)

S6 Fig. Pair-wise BLASTN of three *Echinochloa* transcriptomes.
(PDF)

S7 Fig. Contig number for significantly up-regulated DEGs based on pair-wise comparisons.

(PDF)

S8 Fig. (A) Hierarchical clustering of DEG belonging to CDPKs. (B) Hierarchical clustering of DEG belonging to previously known *Arabidopsis* abiotic stress responsive genes.

(PDF)

S9 Fig. Phylogenetic relationships among three *E. crus-galli* accessions in comparison with *E. oryzicola*, rice (*Oryza sativa*), and sorghum (*Sorghum bicolor*); (A) Phylogenetic relationship using *matK* genes and (B) Phylogenetic relationship using *rbcl* genes.

(PDF)

S1 Table. Primers used for qRT-PCR validation of selected DEGs.

(DOCX)

S2 Table. GO categorization for molecular process using three *E. crus-galli* transcriptomes.

(DOCX)

S3 Table. GO categorization for cellular process using three *E. crus-galli* transcriptomes.

(DOCX)

S4 Table. GO categorization for biological process using three *E. crus-galli* transcriptomes.

(DOCX)

S5 Table. Hierarchical clustering of DEGs belonging to serine/threonine protein kinases (STKs).

(DOCX)

S6 Table. Hierarchical clustering of DEGs belonging to leucine-rich repeat protein kinase (LRR-STKs).

(DOCX)

S7 Table. Hierarchical clustering of DEGs belonging to calcium-related signaling.

(DOCX)

S8 Table. Hierarchical clustering of DEGs belonging to signaling-related proteins.

(DOCX)

S9 Table. Hierarchical clustering of DEGs belonging to glycoproteins.

(DOCX)

S10 Table. Hierarchical clustering of DEGs belonging to glycosidases.

(DOCX)

S11 Table. Hierarchical clustering of DEGs belonging to previously known *Arabidopsis* abiotic stress-responsive genes.

(DOCX)

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Author Contributions

Conceived and designed the experiments: DK GN. Performed the experiments: JI HP. Analyzed the data: GN. Contributed reagents/materials/analysis tools: JK. Wrote the paper: GN DK MY AF TY.

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